

CATEGORY:

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U.S. DEPARTMENT OF CONGUERCE PATENT AND TRADEMARK OFFICE FORM PTO-1390 (REV. 1-98) BO 41497 TRANSMITTAL LETTER TO THE UNITED STATES U.S. APPLICATION NO. (If known, see 37 CFR 1.5 DESIGNATED/ELECTED OFFICE (DO/EO/US) CONCERNING A FILING UNDER 35 U.S.C. 371 PRIORITY DATE INTERNATIONAL FILING DATE INTERNATIONAL APPLICATION NO. 3 April 1998 PCT/NL98/00186 TITLE OF INVENTION METHOD OF INTERSTRAIN DIFFERENTIATION OF BACTERIA Johannes Dirk Anthonie VAN EMBDEN, APPLICANT(S) FOR DO/EO/US Leendert Marinus SCHOULS and Rudolph JANSEN Applicant herewith submits to the United States Designated/Elected Office (DO/EO/US) the following items and other information: 1. X This is a FIRST submission of items concerning a filing under 35 U.S.C. 371. 2. This is a SECOND or SUBSEQUENT submission of items concerning a filing under 35 U.S.C. 371. 3. X This express request to begin national examination procedures (35 U.S.C. 371(f)) at any time rather than delay examination until the expiration of the applicable time limit set in 35 U.S.C. 371(b) and PCT Articles 22 and 39(1). 4. X A proper Demand for International Preliminary Examination was made by the 19th month from the earliest claimed priority date. 3. X A copy of the International Application as filed (35 U.S.C. 371(c)(2)) is transmitted herewith (required only if not transmitted by the International Bureau). has been transmitted by the International Bureau. is not required, as the application was filed in the United States Receiving Office (RO/US). ا ا A translation of the International Application into English (35 U.S.C. 371(c)(2)). Amendments to the claims of the International Aplication under PCT Article 19 (35 U.S.C. 371(c)(3)) are transmitted herewith (required only if not transmitted by the International Bureau). have been transmitted by the International Bureau. have not been made; however, the time limit for making such amendments has NOT expired. have not been made and will not be made. A translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371 (c)(3)). An oath or declaration of the inventor(s) (35 U.S.C. 371(c)(4)). 10. A translation of the annexes of the International Preliminary Examination Report under PCT Article 36 (35 U.S.C. 371(c)(5)). Items 11. to 16. below concern document(s) or information included: 11. An Information Disclosure Statement under 37 CFR 1.97 and 1.98. 12. An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is included. 13. X A FIRST preliminary amendment. A SECOND or SUBSEQUENT proliminary amendment 14. A substitute specification. 15. A change of power of attorney and/or address letter. 16. X Other items or information: International Preliminary Examination Report. Application Data Sheet. Search Report.

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PATENTS

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re application of

Johannes D.A. VAN EMBDEN et al.

Serial No. (unknown)

Filed herewith

METHOD INTERSTRAIN DIFFERENTIATION OF BACTERIA

PRELIMINARY AMENDMENT

Commissioner for Patents

Washington, D.C. 20231

Sir:

Prior to calculation of the filing fee, please amend the above-identified application as follows:

IN THE DRAWINGS:

Please add Figure 6 as filed in the Article 34 amendment of June 2, 2000. The sheet containing Figure 6 is attached hereto and marked "AMENDED SHEET".

IN THE CLAIMS:

Please substitute Claims 1-25 as originally filed, which appear on pages 22-27, with Claims 1-25 as filed in the Article 34 amendment of June 15, 2000. The pages containing Claims 1-25 are marked "AMENDED SHEET" and are attached hereto. Following the insertion of Claims 1-25, please amend these claims as follows:

Claim 3, line 1, cancel "or 2".

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Claim 4, line 1, change "any of the preceding claims" to --claim 1--.

Claim 5, line 1, change "any of the preceding claims" to --claim 1--.

Claim 6, line 1, change "any of the preceding claims" to --claim 1--.

Claim 7, line 1, change "any of the preceding claims" to --claim 1--.

Claim 8, line 1, change "any of the preceding claims" to --claim 1--.

Claim 10, line 1, change "any of the preceding claims" to --claim 1--.

Claim 11, line 1, change "any of the preceding claims" to --claim 1--.

Claim 14, line 1, cancel "or 13".

Claim 15, line 1, change "any of claims 12-14" to --claim 12--.

Claim 16, line 1, change "any of claims 12-15" to --claim 12--.

Claim 17, line 3, change "any of claims 12-16" to --claim 12--.

Claim 19, line 1, cancel "or 18".

Claim 21, line 1, change "claim 21" to --claim 20--.

Amend Claim 24 as follows:

--24. (amended) Kit for carrying out a method [according to any of claims 1-19] of in vitro amplification of nucleic acid using amplification primers, comprising a primer pair according to [any of claims 20-23] claim 20 and an

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oligonucleotide probe or a carrier, said carrier comprising at least 1 oligonucleotide probe specific for a spacer region of a bacterium to be determined said bacterium not belonging to M. tuberculosis complex, said oligonucleotide probe being an oligonucleotide probe comprising between 12 to 40 nucleotides, said probe being sufficiently homologous to any of the spacer sequences or to fragments or derivatives of such spacer sequences to hybridise to such a spacer sequence, said oligonucleotide probe comprising at least ten consecutive nucleotides homologous to such a spacer sequence and/or exhibiting at least 60% homology, with the corresponding part of the spacer sequence.--

REMARKS

The above changes in the drawings and claims merely place the national phase application in the same condition as it was during Chapter II cf the international phase, with the multiple dependencies in the claims being removed.

Respectfully submitted,

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Ву

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October 3, 2000

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Claims

- 1. A method of in vitro amplification of nucleic acid using amplification primers in a manner known per se, in amplification reactions such as PCR, LCR or NASBA, wherein a pair of primers is used comprising oligonucleotide sequences sufficiently complementary to a part of the Direct Repeat sequence of a bacterium other than a microorganism belonging to the Mycobacterium tuberculosis complex of microorganisms for hybridisation to a Direct Repeat to occur and subsequently elongation of the hybridised primer to take place, said primers being such that elongation in the amplification reaction occurs for one primer in the 5' Direction and for the other primer in the 3' Direction, wherein the Direct Repeat is a sequence with a length between 20-50 base pairs which occurs 5-60 times in a region of the bacterial genome, whereby the Direct Repeat sequences are separated by spacer sequences with a length of between 20-50 nucleotides, said spacer sequences being non repetitive.
 - 2. A method according to claim 1 wherein the Direct Repeat sequence is obtainable from screening a genomic bacterial nucleic acid sequence using the programme Patscan wherein the Direct Repeat is designated p1 with a length between 20-50 basepairs then p1 is sought 20-50 basepairs downstream of p1 as the pattern

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pI=(20..50)(20..50)p1(20..50)p1 or a variant thereof wherein the ranges of the nucleotide lengths are shorter and wherein the frequency of occurrence of the Direct Repeat can vary between 5 and 60.

- 5 3. A method according to claim 1 or 2 wherein the Direct Repeat has a length between 30-40 base pairs and the spacer has a length of 35-45 base pairs.
 - 4. A method according to any of the preceding claims wherein the Direct Repeat has a terminus exhibiting at least 3 out of 5 nucleotides identical with the sequence GAAAC, preferably 4, said termini being selected from GAAAC, GAAXXC GAACTC, GXAAC, GCAAC, GAAA, GAAXC, GAAGC, AAAC.
 - 5. A method according to any of the preceding claims wherein the Direct Repeat terminates with GAACTC, ATACAC, AAAACT, TTGCAA, GGAAAC, TGAAAC, TGAAAC, TGGAAA, TTTAAC, TGAAAT or TTCAAC.
 - 6. A method according to any of the preceding claims wherein the Direct Repeat has stretches of 3-4 identical bases.
- 7. A method according to any of the preceding claims wherein the Direct Repeat has a sequence such that it is not prone to loop formation or any other obvious secondary structure.
- 8. A method according to any of the preceding claims wherein the bacterium is 25 a pathogenic bacterium selected from the group of Gram negative aerobic/microaerophilic rods and cocci and facultatively anaerobic gram-negative rods.
- A method according to claim 8 wherein the bacterium is a pathogenic bacterium selected from the genera Escherichia, Shigella, Salmonella, Klebsiella,
 Enterobacter, Yersinia, Serratia, Haemophilus, Vibrio, Legionella, Neisseria,
 Pseudomonas, Bordetella, Staphylococcus, Streptococcus and Acinetobacter.

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- A method according to any of the preceding claims, wherein said primers 10. have oligonucleotide sequences complementary to non overlapping parts of the Direct Repeat sequence and such that the elongation reactions from each primer can occur without hindrance of the other when both primers hybridise to the same Direct Repeat and undergo elongation.
- A method according to any of the preceding claims, wherein one primer 11. DRa is capable of elongation in the 5' Direction and the other primer DRb is capable of elongation in the 3' Direction and DRa is complementary to a sequence of the Direct Repeat located to the 5' side of the sequence of the Direct Repeat to which DRb is complementary.
- A method of detection of a bacterium, said bacterium not belonging to the M 12. tuberculosis complex of microorganisms said method comprising
- amplifying nucleic acid from a sample with the method according to any of the 1) preceding claims, followed by
- carrying out a hybridisation test in a manner known per se, wherein the 2) amplification product is hybridised to an oligonucleotide probe or a plurality of different oligonucleotide probes, each oligonucleotide being sufficiently homologous to a part of a spacer of the Direct Region of the bacterium to be determined for hybridisation to occur to amplified product if such spacer nucleic acid was present in the sample prior to amplification, said hybridisation step optionally being carried out without prior electrophoresis or separation of the amplified product.
- 25 detecting any hybridised products in a manner known per se. 3)
 - A method according to claim 12, wherein the hybridisation test is carried out 13. using a number of oligonucleotide probes, said number comprising at least a number of oligonucleotides probes specific for the total spectrum of bacteria it is desired to detect.

- A method according to claim 12 or 13, wherein the oligonucleotide probe is at least ten oligonucleotides long and is a sequence complementary to a sequence selected from any of the spacer sequences of the Direct Repeat region of the bacterium to be determined or is a sequence complementary to fragments or derivatives of said spacer sequences, said oligonucleotide probe being capable of hybridising to such a spacer sequence and comprising at least ten consecutive nucleotides homologous to such a spacer sequence and/or exhibiting at least 60% homology, preferably exhibiting at least 80% homology with such a spacer sequence.
- 15. A method according to any of claims 12-14 wherein the bacterium is a pathogenic bacterium selected from the group of Gram negative aerobic/microaerophilic rods and cocci and facultatively anaerobic Gram-negative rods.
- 16. A method according to any of claims 12-15 wherein the bacterium is a pathogenic bacterium selected from the genera Escherichia, Shigella, Salmonella, Klebsiella, Enterobacter, Yersinia, Serratia, Haemophilus, Vibrio, Legionella, Neisseria, Pseudomonas and Bordetella and the group of Gram positive bacterial genera Staphylococcus and Streptococcus as target for the differentiation method.
- 20 17. A method for differentiating the type of bacterium in a sample, said bacterium not belonging to the M. tuberculosis complex, said method comprising carrying out the method according to any of claims 12-16, followed by comparison of the hybridisation pattern obtained with a reference.
- 25 18. A method according to claim 17, wherein the reference is the hybridisation pattern obtained with one or more known strains of the bacterium to be determined in analogous manner.
- 19. A method according to claim 17 or 18 wherein the reference is a source providing a list of spacer sequences and sources thereof, such as a data bank.

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- 20. A pair of primers wherein both primers comprise oligonucleotide sequences of at least 7 oligonucleotides and are sufficiently complementary to a part of the Direct Repeat sequence of the microorganism E. coli for hybridisation to occur and subsequently elongation of the hybridised primer to take place, said primers being such that elongation in the amplification reaction occurs for one primer in the 5' Direction and for the other primer in the 3' Direction and wherein sufficiently complementary means said oligonucleotide sequence comprises at least seven consecutive nucleotides homologous to such a Direct Repeat sequence in particular a Sequence described inSEQ ID No. 1-18 and/or exhibits at least 60% homology, with the corresponding part of the Direct Repeat sequence.
- 21. Primer pair according to claim 21, comprising one primer DRa capable of elongation in the 5' Direction and the other primer DRb capable of elongation in the 3' Direction with DRa being complementary to a sequence of the Direct Repeat located to the 5' side of the sequence of the Direct Repeat to which DRb is complementary, the Direct Repeat being present in the Direct Region of SEQ IS No. 2.
- 22. A pair of primers wherein both primers comprise oligonucleotide sequences of at least 7 oligonucleotides and are sufficiently complementary to a part of the Direct 20 Repeat sequence of the microorganism S. typhimurium for hybridisation to occur and subsequently elongation of the hybridised primer to take place, said primers being such that elongation in the amplification reaction occurs for one primer in the 5' Direction and for the other primer in the 3' Direction and wherein sufficiently complementary means said oligonucleotide sequence comprises at least seven consecutive nucleotides homologous to such a Direct Repeat sequence in particular the DEQ ID No. 2 and/or exhibits at least 60% homology, with the corresponding part of the Direct Repeat sequence.
- 23. Primer pair according to claim 22, comprising one primer DRa capable of elongation in the 5' Direction and the other primer DRb capable of elongation in the 3' 30 Direction with DRa being complementary to a sequence of the Direct Repeat located to the 5' side of the sequence of the Direct Repeat to which DRb is complementary, the Direct Repeat being present in the Direct Region of SEQ ID No. 8.

AMENDED SHEET

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- 24. Kit for carrying out a method according to any of claims 1-19, comprising a primer pair according to any of claims 20-23 and an oligonucleotide probe or a carrier, said carrier comprising at least 1 oligonucleotide probe specific for a spacer region of a bacterium to be determined said bacterium not belonging to M. tuberculosis complex, said oligonucleotide probe being an oligonucleotide probe comprising between 12 to 40 nucleotides, said probe being sufficiently homologous to any of the spacer sequences or to fragments or derivatives of such spacer sequences to hybridise to such a spacer sequence, said oligonucleotide probe comprising at least ten consecutive nucleotides homologous to such a spacer sequence and/or exhibiting at least 60% homology, with the corresponding part of the spacer sequence.
- 25. Kit according to claim 24 further comprising a data carrier with required reference patterns of the bacterial strain to be determined.

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A method of interstrain differentiation of bacteria.

Summary of the invention

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The subject invention lies in the field of interstrain differentiation of bacteria. A general method has been developed with which various types of bacteria can be differentiated into separate individual strains. Thus in particular in the clinical setting this method can suitably be used to determine what strain of bacterium is present in a sample. This new method is applicable for discerning between various strains of both Gram negative and Gram positive types of bacteria.

Background of the invention

Previously we had disclosed a method called oligotyping for interstrain differentiation of Mycobacterium tuberculosis strains in W095/31569. It was stated in this document that one of the key factors in the control of tuberculosis is the rapid diagnosis of the disease and the identification of the sources of infection. M. tuberculosis strain typing has already proved to be extremely useful in outbreak investigations (6, 14, 31) and is being applied to a variety of epidemiologic questions in numerous laboratories. Traditionally, laboratory diagnosis is done by microscopy, culturing of the micro-organism, skin testing and X-ray imaging. Unfortunately, these methods are often not sensitive, not specific and are very time-consuming, due to the slow growth rate of M. tuberculosis. Therefore, new techniques like in vitro amplification of M. tuberculosis DNA have been developed to rapidly detect the micro-organism in clinical specimens (14). The ability to differentiate isolates of M. tuberculosis by DNA techniques has revolutionarized the potential to identify the sources of infection and to establish main routes of transmission and risk factors for acquiring tuberculosis by infection (1,3-10, 14, 16, 19-22, 25, 26, 27-33). The use of an effective universal typing system will allow strains from different geographic areas to be compared and the movement of individual strains to be tracked. Such data may provide important insights and identify strains with particular problems such as high infectivity, high virulence and/or multidrug resistance. Analysis of large numbers of isolates may provide answers to long-standing questions regarding the efficacy of BCG vaccination and the

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frequency of reactivation versus reinfection.

The same problems identified for M. tuberculosis are inherent in differentiation of numerous other bacteria. The problems specifically arise for potentially epidemic pathogens and for bacteria that infect hospitals. A more rapid and simple typing method is required. Preferably the testing methods for various bacteria will occur in the same manner ensuring routine use for all types of bacteria for which testing is required. Preferably a test that can be carried out by non specialised personnel using little laboratory space and time is sought after.

The method disclosed in W095/31569 is based on the DNA polymorphism found at a unique chromosomal locus, the "Direct Repeat" (DR) region, which is uniquely present in M. tuberculosis complex bacteria. This locus was discovered by Hermans et al. (15) in M. bovis BCG, the strain used worldwide to vaccinate against tuberculosis. The DR region in M. bovis BCG consists of Directly repeated sequences of 36 base pairs, which are interspersed by non-repetitive DNA spacers, each 35 to 41 base pairs in length (15). The number of copies of the DR sequence in M.bovis BCG was determined to be 49. In other strains of the M. tuberculosis complex the number of DR elements was found to vary (15). The vast majority of the M. tuberculosis strains contain one or more IS6110 elements in the DR containing region of the genome.

It has been shown (12) that the genetic diversity in the DR region is generated by differences in the DR copy number, suggesting that homologeous recombination between DR sequences may be a major driving force for the DR-associated DNA polymorphism (12). The high degree of DNA polymorphism within a relatively small part of the chromosome makes this region well-suited for a PCR-based fingerprinting technique.

Figure 1 depicts the structure of the DR region of M. bovis BCG as determined previously by Hermans et al. and Groenen et al. (12, 15). For the sake of convenience we will designate a DR plus its 3'adjacent spacer sequence as a "Direct Variant Repeat" (DVR). Thus, the DR region is composed of a discrete number of DVR's, each consisting of a constant part (DR) and a variable part (the spacer).

The method disclosed in W095/31569 is based on a unique method of in vitro amplification of DNA sequences within the DR region and the hybridisation of the amplified DNA with multiple, short synthetic oligomeric DNA sequences based on the sequences of the unique spacer DNA's within the DR region (figure 2). This differs from previous PCR methods in the use of a set of primers with both primers having multiple

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priming sites as opposed to having one of the primers bind to a fixed priming site such as to a part of IS6110. Because M. tuberculosis complex strains differ in the presence of these spacer sequences, strains can be differentiated by the different hybridisation patterns with a set of various spacer DNA sequences.

The method consists of in vitro amplification of nucleic acid using amplification primers in a manner known per se in amplification reactions such as PCR, LCR or NASBA, wherein a pair of primers is used comprising oligonucleotide sequences sufficiently complementary to a part of the Direct Repeat sequence of a microorganism belonging to the M. tuberculosis complex of microorganisms for hybridisation to a Direct Repeat to occur and subsequently elongation of the hybridized primer to take place, said primer being such that elongation in the amplification reaction occurs for one primer in the 5' Direction and for the other primer in the 3' Direction. Due to the multiple presence of Direct Repeats in the microorganisms to be detected the use of such primers implies that all the spacer regions will be amplified in an efficient manner. In particular it is not necessary for extremely long sequences to be produced in order to obtain amplification of spacers located at a distance from the primer. With the instant selection of the primer pairs a heterogenous product is obtained comprising fragments all comprising spacer region nucleic acid. Subsequently the detection of the amplified product can occur simply by using an oligonucleotide probe directed at one or more of the spacer regions one wishes to detect. In order to avoid hindrance in the amplification reactions the primers can oligonucleotide sequences complementary to non-overlapping parts of the Direct Repeat sequence so that when both primers hybridize to the same Direct Repeat and undergo elongation they will not be hindered by each other. In particular to avoid any hindrance during elongation reactions when one primer DRa is capable of elongation in the 5' Direction and the other primer DRb is capable of elongation in the 3' Direction the DRa is selected such that it is complementary to a sequence of the Direct Repeat located to the 5' side of the sequence of the Direct Repeat to which DRb is complementary. The primer used must have an oligonucleotide sequence capable of annealing to the consensus sequence of the Direct Repeat in a manner sufficient for amplification to occur under the circumstances of the particular amplification reaction. A person skilled in the art of amplification reactions will have no difficulty in determining which length and which degree of homology is required for good amplification

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reactions to occur. The consensus sequence of the Direct Repeat of microorganisms belonging to the M. tuberculosis complex is given in sequence id. no. 2 and in figure 1.

In addition to what has already been disclosed in W095/31569 we also determined the spoligotypes of M. tuberculosis strains which were subcultured for many months both in the laboratory and in guinea pigs. The strains selected for this purpose were those used in a previous study on the stability of TS 6110 (2). All subcultured strains displayed the identical spoligotype patterns compared with the primary cultures thus indicating the pace of the molecular clock in this instance is slow enough for use in epidemiology of the disease.

Because of the large success and simplicity of the method for Mycobacterium tuberculosis strain differentiation and in view of problems in strain differentiation with other microorganisms we used the Direct Repeat consensus sequence to screen data bases with nucleic acid sequences from other microorganisms. Unfortunately no further matches were found. The Direct Repeat sequence appeared to be unique for the Mycobacterium tuberculosis as did their spacer sequences. As to date no function had actually been attributed to the Direct Repeat sequence it was unexpected that the sequence was universally distributed amoung other types of microorganisms. Such would at best be expected if the sequence had a function that was required also in other organisms.

Description of the invention

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Notwithstanding the negative result after screening with the Direct Repeat consensus sequence we considered further analysis of known sequences by looking for a pattern in the nucleic acid sequences of other microorganisms reminiscent of the Direct Repeat-spacer pattern in Mycobacterium tuberculosis. Quite unexpectedly we found using a specifically designed computer programme that such patterns existed in a large number of other microorganisms with a broad range of genera. It appears that the DR-like sequences are very common in prokaryotes. They are however noticeably absent in eukaryotes. Chapter III of Bergeys Determinative Manual of Bacteriology Ninth edition (11) provides a table of characteristics for distinguishing prokaryotes from eukaryotes i.e. distinguish bacterium from microoscopic eukaryotes in the shape of mold, yeast, algae or protozoans.

All bacterial sequences analysed revealed the presence of such

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a sequence structure and thus the oligotyping method illustrated for Mycobacterium tuberculosis can be applied for differentiating between all strains of bacteria. It was totally unexpected that a consensus structure of this type could be universally found. The Direct Repeat sequences themselves are different between different genera but the general framework of a cluster of Direct Repeat sequences, separated by a number of non repetitive spacers is universally present in bacterial genomes. Considering the fact that thusfar no function has been attributed to such a region in Mycobacterium tuberculosis or in fact for any of the sequences comprising Direct Repeat like regions in any other bacteria for which such sequences had been described this is remarkable.

Bacteria can be divided into Archaebacteria and Eubacteria. The eubacteria in turn can de distinguished into Gram-negative and Grampositive bacteria with cell walls and Eubacteria lacking cell walls. Chapter IV of Bergeys determinative Manual of Bacteriology Ninth edition (11) reveals the characteristics for each group. Over a wide range of the subgroups in these 4 groups we have found the presence of the consensus structure i.e. the presence of DR-like loci. The IV groups have been subdivided by Bergey into more than 30 subgroups. We have examples in Groups 3,4,5 and 6, Group 11, 17, 31, 32, 33. The method according to the invention is particularly of interest for the bacteria that are pathogenic for humans. Group 4 comprises Gram negative bacteria. Genera from Group 4 are Legionella (which can cause pneumonia) and Legionnaires disease, the genus Neisseria (of which Neisseria meningitidis is well known as causative agent of meningitis and of which Neisseria gonorrhoeae is another example). the genus Pseudomonas (renown for hospital infections) and the genus Bordetella (of which Bordetella pertussis is well known as causative agent of whooping cough). In Group 5 bacteria as defined in Bergeys Manual the Enterobacteriacae form a family of 30 genera. These bacteria form a particularly interesting group of Gram negative bacteria that infect humans. Suitable examples of genera from family are Enterobacter, Escherichia, Shigella, Salmonella, Serratia, Klebsiella and Yersinia. Other less well known pathogenic Enterobacteriacae genera are Cedeca, Citrobacter, Kluyvera, Leclercia, Pantoea, Proteus, Providencia and Hafnia. Other Group 5 families are Pasteurellaceae with the genus Haemophilus and the family Vibrionaceae with the genus Vibrio. Haemophilus influenzae is a leading cause of meningitis in children and also other septicemia conditions. Vibrio cholerae is the causative agent of cholera, V. parahaemolyticus can cause

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food poisoning and V. vulnificus causes highly fatal septicemia.

Of the Enterobacteriacae Shigella, Escherichia and Salmonella are best known and difficult to differentiate. Shigella is an intestinal pathogen of humans causing bacillary dysentery. Well known strains are S. dysenteriae, S. flexneri, S. boydii, S. sonnei. The genus Salmonella is well known for food poisoning. Well known Salmonella strains are S. typhimurium, S. arizona, S. choleraesuis, S. bongori. Salmonella are also causative agents of typhoid fever, enteric fevers, gastroenteritis and septicemia. The genus Serratia bacteria are opportunistic pathogens for hospitalized humans causing septicemia and urinary tract infections. Examples are S. liquefaciens and S. marcescens. Of the Escherichia E. coli is best known as major cause of urinary tract infections and nosocomial infections including septicemia and meningitis. Other species are usually associated with wound infections.

Enterobacter constitutes a problem genus of opportunistic pathogens causing burn wound and urinary tract infections occasionally also meningitis and septicemia. Well known species are E. cloacae, E. sakazakii, E. aerogenes, E. agglomerans, E. gergoviae. Klebsiella are also causative agents of bacteriemia, pneumonia, urinary tact and other human infections in urological, neonatal, intensive care and geriatric patients. Klebsiella pneumoniae and K. oxytoca are examples of species in the genus.

Particularly interesting from a clinical point of view are also the Gram positive pathogenic bacteria. The genera Streptococcus and Staphylococcus form examples of such bacteria. Streptococcus pneumoniae, Streptococcus pyogenes and Staphylococcus aureus are examples thereof. Of the mentioned groups and genera the pathogenic bacteria are of interest. These bacteria are dangerous when infecting hospitals in particular.

Due to the increasing incidence of infection differentiation of potentially epidemiological organisms is also of interest. Such organisms comprise Bordetella pertussis and Neisseria menigitidis the causative organism of meningitis is of particular interest. Quite specifically pathogenic bacteria infecting hospitals and bacteria capable of causing epidemics are targets for the differentiation method according to the invention.

The invention consists of a method of in vitro amplification of nucleic acid using amplification primers in a manner known per se, in amplification reactions such as PCR, LCR or NASBA, wherein a pair of primers is used comprising oligonucleotide sequences sufficiently

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complementary to a part of the Direct Repeat sequence of a bacterium other than a microorganism belonging to the M tuberculosis complex of microorganisms for hybridisation to a Direct Repeat to occur and subsequently elongation of the hybridised primer to take place, said primers being such that elongation in the amplification reaction occurs for one primer in the 5' Direction and for the other primer in the 3' Direction, wherein the Direct Repeat is a sequence with a length between 20-50 base pairs which occurs 5-60 times in a contiguous region of the bacterial genome, whereby the Direct Repeat sequences are separated by spacer sequences with a length of between 20-50 nucleotides, said spacer sequences being non repetitive. By using the programme Patscan e.g. on the nucleic acid data bases for microorganism genomic sequences such motifs and thus also the identities of the various species specific Direct Repeats and the corresponding spacer sequences can be obtained. In the Patscan programme the Direct Repeat can be designated p1 with a length between 20-50 basepairs then search for p1 20-50 basepairs downstream of pl. Thus this pattern in Patscan is described p1=(20..50)(20..50)p1(20..50)p1. The length of the sequences can be varied as can the intermediate distance and the number of times the Direct Repeat has to occur. A Direct Repeat can often have a length of 30-40 base pairs with a spacer length of 35-45 base pairs. Basically we looked for a stretch of identical repeat sequences interspersed by spacer sequences which do not necessarily share much of their sequence with the Direct Repeat of M. tuberculosis. The patscan programme is freely accessible at the Internet site: http://www-c.mcs.anl.gov/home/overbeek/-PatScan/HTML/patscan.html. The programme was written by Ross Overbeek Mathmatics and Computer Science Division Argonne National Laboratory Building 221 Room D-236 9700 S. Cass Avenue Arginne IL 60439 USA.

Most of the Repeats exhibit one or more of the following characteristics, they end with a sequence similar to GAAAC i.e. exhibit at least 3 of the nucleotides of this consensus sequence at the terminus, preferably 4 or 5, start with CTTTG, have stretches of 3-4 identical bases. The termini can for example be selected from GAAAC, GAAXXC GAACTC, GXAAC, GCAAC, GAAA, GAAXC, GAAGC and AAAC. Suitable Termini are provided in Table II.

Organisms as diverse as the Archaebacteria e.g. Methanococcus jannasschi (Group 31), Haloferax mediterranei (Group 33), the cyanobacteria Calotrix (Group 11), and Anabeana (Group 11), and purple bacteria e.g. E.coli (Group 5), Mycobacterium tuberculosis (Group 21) and

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Thermus thermophilus (Group 4), Archaeoglobus (Group 32) and Thermotoga (Group 6) were found to possess DR-like sequences upon analysis of their genomes using the Patscan programme. In the subsequent study of literature from which these data were derived it also became clear from Southern blots that the Repeat sequences were also found in related species.

With regard to the genetic organisation the structures of the DR-like loci in the microorganisms is rather variable (figure 3). In M. tuberculosis the DR locus is large and in most isolates it is disrupted by an insertion element. This is also the case in T. thermophilus, however here the number of DVR's is only 11 and the DR locus is disrupted by two insertion elements. In E. coli K12 2 DR loci are present separated by approximately 22kb; in Anabaena the locus is of intermediate size and interrupted by a 130 bp sequence of unknown function or origin. In H. mediterranei the DR locus is of intermediate size and not disrupted, however there is evidence for a second DR locus on one of the mega plasmids found in this organism. In M. jannaschii there is one locus of intermediate size but at several other positions in the genome one or a few other DVR's are found. In most cases the DVR's are linked to a socalled Long Repeat (LR) element of unknown function. Also in M. jannaschii mega plasmids are found but in contrast to H. mediterranei they do not contain DR sequences.

Accession numbers for the sequences of various organisms for which the DR like loc1 have been found are provided here. For E. coli and Shigella M27059, M27060, U29579, U29580 and M18270. The relevant portions of the sequences are also disclosed by Blattner for E. coli. Nakata et al reveal in the Journal of Bacteriology (13) that downstream of the iap region a sequence of 29 bases appears 14 times 32 or 33 base pairs apart. Nucleotide sequences hybridizing to the 29 base pair sequence were also detected in Shigella dysenteriae and Salmonella typhimurium.

A DR-like sequence was found in the contig 214 of S. pyogenes M1(ATCC 700294) of the genome sequencing project of the University of Oklahoma. Further research into this DR-like sequence in other S. pyogenes revealed spacer polymorphism. The DR regions of eight S. pyogenes isolates were studied. The DR regions were isolated by PCR using primers that were derived from the database (University of Oklahoma, serotype M1 ATCC 700294. The sequence data is available under http://www.genome.ou.edu. This strain contains seven repeats and six spacers.

Five of the isolates gave a PCR product, these were a M2 strain, a M4 strain and three M1 strains. The M4 strain contained only a single repeat sequence that was flanked by the same sequences as the ATCC 700294. The M2 strain sequencing did not work, but the size of the PCR fragment indicated that two repeats are present. The three M1 strains were all the same, they contained four repeats and three spacers. The repeats were identical to ATCC 700294, while one of the spacers was identical to ATCC 700294 and two were different.

These studies on *S. pyogenes* show that the DR regions have conserved spacers and repeat sequences.

The Salmonella genomic sequence as sequenced by the University of Washington St Louis has also revealed the presence of DR-like sequences. The DR exhibits high homology with the Direct Repeat of E. coli. One of the contigs revealed 7 Repeats and 6 spacers.

A panel of five $E.\ coli$ isolates and three Shigella strains were studied. The five $E.\ coli$ isolates were selected to have an optimal diversity, they were isolated from different species or geographic regions. The Shigella strains are considered separate (sub)species. See Table 1. The isolates were obtained from the collection of Dr. Wim Gaastra.

Table 1

species	descript:on	DRI*	DRII*
E. coli 184	American isolate	Southern	PCR
358	human urinary tract	Southern	Southern
968	mastitis	Southern	PCR
1008	chicken	PCR	PCR
1732	human intestine	Southern	PCR
Shigella disenteriae	593	Southern	PCR
sonnei	595	Southern	PCr
boydil	603	Southern	PCR

* The DR regions were identified by Southern blot of genomic DNA and DRI and DRII regions of *E. coli* K12. When PCR is indicated the DR regions were identified by the Southern and the PCR. This PCR was done with primers derived from the K12 sequence.

The DRI and DRII sequences that could be amplified by PCR were

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cloned and sequenced. Somehow the DRI regions could not be amplified by PCR using the primers designed on the K12 sequence, while the Southern data demonstrate that DRI is present. Apparently, the recognitions sites for the primers are polymorphic. The sizes of the DRII regions were found to vary greatly between these isolates. The smallest was a single repeat in the S. sonnei strain and the largest was a repeat cluster of at least 15 repeats in E. coli isolate 1008. The sequences of the repeats were highly conserved between these isolates. The S. typhimurium data is obtainable from the Internet http://genome.wustl.edu/gsc/bacterial/salmonella.html.

The spacer sequences almost all were unique. Approximately 40 spacers have been sequenced and only three of them were already known from a previously sequenced DR region. This indicates a high number of different spacer sequences in *E. coli*.

Accession number X73453 provides the Halerofax mediterranei sequence. The sequence can also be found in Molecular Microbiology 17 of 1995 in an article by Mojica et al. (17). The Repeat sequence has also been found in related species.

The genomic project of the Methanococcus jannaschii reveals a DR-like sequence as is apparent from the Bult et al article in Science 273 of 1996 (18). The Accession number is U67459 i.a.

Accession number X87270 for Anabeana sp reveals 17 spacers and a LTRR element. These elements also occur in related species of cyanobacteria such as Calotrix. The sequence data are provided by Masepohl et al in BBA 1307 1996 (23).

Accession number AE000782 for Archaeoglobus fulgidus reveals three DR-like Repeats with the same Repeat sequence and the this has a slightly larger but closely related Repeat. The Repeats are present 20-30 times. The spacers are unique sequences. H.P. Klenk discloses sequence data in Nature 390 1997 (24).

The invention also covers a method of detection of a bacterium, said bacterium not belonging to the M. tuberculosis complex of microorganisms said method comprising

- 1) amplifying nucleic acid from a sample with the amplification method according to any of the preceding described embodiments of the amplification method according to the invention, followed by
- 2) carrying out a hybridisation test in a manner known per se, wherein the amplification product is hybridised to an oligonucleotide probe or a plurality of different oligonucleotide probes, each

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oligonucleotide being sufficiently homologous to a part of a spacer of the Direct Region of the bacterium to be determined for hybridisation to occur to amplified product if such spacer nucleic acid was present in the sample prior to amplification, said hybridisation step optionally being carried out without prior electrophoresis or separation of the amplified product.

3) detecting any hybridised products in a manner known per se.

The method can be carried out in a manner such that the hybridisation test is carried out using a number of oligonucleotide probes, said number comprising at least a number of oligonucleotides probes specific for the total spectrum of bacteria it is desired to detect. In a suitable embodiment of a method according to the invention the oligonucleotide probe is at least seven oligonucleotides long and is a sequence complementary to a sequence selected from any of the spacer sequences of the Direct Repeat region of the bacterium to be determined or is a sequence complementary to fragments or derivatives of said spacer sequences, said oligonucleotide probe being capable of hybridising to such a spacer sequence and comprising at least seven consecutive nucleotides homologous to such a spacer sequence and/or exhibiting at least 60% homology, preferably exhibiting at least 80% homology with such a spacer sequence.

Preferably the method according to the invention is carried out to determine the presence and nature of a pathogenic bacterium selected from the group of Gram negative bacteria of Groups 4 and 5 of Bergeys Determinative Manual of Bacteriology ninth edition. Of particular interest due to damage caused by such pathogens are bacteria belonging to the families Enterobacteriaceae, Pasteurellaceae and Vibrionaceae of Group 5, most specifically the Enterobacteriaceae. Also of interest are the Gram positive bacteria of Group 17. Suitable examples of genera of the pathogenic bacterium to be detected from the group of Gram negative bacteria of Bergeys Determinative Manual of Bacteriology ninth edition Eschericchia, Shigella, Salmonella, Klebsiella, Enterobacter, Yersinia. Serratia, Haemophilus, Vibrio, Legionella, Neisseria, Pseudomonas and Bordetella. For the group of Gram positive bacterial genera Staphylococcus and Streptococcus are targets differentiation method.

Suitably in a method according to the invention for differentiating the type of bacterium in a sample, said bacterium not belonging to the M. tuberculosis complex the hybridisation pattern is

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compared with that obtained with a reference. Such a reference can be the hybridisation pattern obtained with one or more known strains of the bacterium to be determined in analogous manner as the strain to be determined. Alternatively the reference is a source providing a list of spacer sequences and sources thereof, such as a data bank. Table II exhibits some suitable examples of sequences that occur as Direct Repeat sequences according to the invention for the genera illustrated.

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Species	Repeat sequence	Reference	EMBL/Genbank
			accession number
Murobacterium tuberculosis	GTCGTCAGACCCAAAACCCCGAGAGGGACGGAAAC	Hermans et al.	
Hospowiebia coli	CGGTTTATCCCCGCTGGCGCGGGGAACTC	Nakata et al.J.Bact.171	M27059 and
200 81101 191007		3553-3556 (1989)	M27060
Shiaella disenteriae	CGGTTTATCCCCGCTGGCGCGGGGAACTC	our own data	
Shige I la some i	CGCITTATCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	our own data	
Shige 11a boudii	CGGITTATCCCCGCTGGCGCGGGGAACTC	our own data	
Salmonella enteritidis	CGGTTTATCCCCGCTGGCGCGGGGAACTC	our own data	
Seratia marcescens	CGGTTTATCCCCGCTGGCGCGGGGAACTC	our own data	
Salmonella tuphimurium	CGGTTTATCCCCGCTGGCGCGGATACAC	contig 70A06 of the	
		typhimurium genome	
		project.Univ. of	
		Washington St. Louis	
Strentococcus puodenes	GTTTTAGAGCTATGCTGTTTTGAATGGTCCCAAAACT	Contig 214 of the S.	
		pyogenes genome project	
		University of Oklahoma	

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Species	Repeat sequence	Reference	EMBL/Genbank
מאַנירוּטמּ			accession number
Thomas aduations	AATCCCCTTACGGGGCTCAATCCCTTGCAA	Ashby et al. Plasmid	M33159
the many inc		24 1-11 (1990)	
Thermotogg maritima	GITTCAATACTTCCTTAGAGGTATGGAAAC	Liebl.et al.	269341
2000 200 200 200 200 200 200 200 200 20		Microbiol.142 2533-	
		2542 (1996)	
Anahaena	GTTTTAACTAACAAAATCCCTATCAGGGATTGAAAC	Masepohl BBA 1307 26-	x87270
		30 1996	
Calotnix	GTITTAAACTITATAAAATCCCTITITAGGGATTGAAAC	idem	Z47161
Haloferax mediterranei	GTTACAGACGAACCCTAGTTGGGTTGAAGC	Mojica et	x7453
		al.Molec.Microbiol.17	
		85-93 (1995)	
Methonocoens ionnaschii	AATTAAAATCAGACCGTTTCGGAATGGAAA	Bult et al.Science	U67459 (for the
2		273 1058-1073 (1996)	large DR-like
			region)
Methomobocterium	ATTTCAATCCCATTTTGGTCTGATTTTAAC	Smith et al. J. Bact.	AE000920 (for R2)
		179 7135-7155 (1997)	
+ hormonitotrophicum	GTTAAAATCAGACCAAAATGGGATTGAAAT		AE000878 (for R1)
Anchocoglobus fulgidus	CTTTCAATCCCATTTTGGTCTGATTTCAAC	Klenk et al. Nature	AE000782 (whole
		390 364-370 (1997)	genome)

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Not only the above methods fall within the scope of the invention but also specifically selected primer pairs for carrying out such a method. A pair of primers according to the invention is a pair wherein both primers comprise oligonucleotide sequences of at least 7 oligonucleotides and are sufficiently complementary to a part of the Direct Repeat sequence of the microorganism E. coli for hybridisation to occur and subsequently elongation of the hybridised primer to take place, said primers being such that elongation in the amplification reaction occurs for one primer in the 5' Direction and for the other primer in the wherein sufficiently 31 Direction and complementary means oligonucleotide sequence comprises at least seven consecutive nucleotides homologous to such a Direct Repeat sequence and/or exhibits at least 60% homology, preferably at least 80% homology, most preferably more than 90% homology with the corresponding part of the Direct Repeat sequence. Suitable Direct Repeat sequences are provided in Table II. In particular such a primer pair can comprise one primer DRa capable of elongation in the 5' Direction and the other primer DRb capable of elongation in the 3' Direction with DRa being complementary to a sequence of the Direct Repeat located to the 5' side of the sequence of the Direct Repeat to which DRb is complementary, the Direct Repeat being present in the Direct Region of E. coli. Another suitable pair comprises primers with oligonucleotide least 7 oligonucleotides sequences ofat and are sufficiently complementary to a part of the Direct Repeat sequence of the microorganism S. typhimurium for hybridisation to occur and subsequently elongation of the hybridised primer to take place, said primers being such that elongation in the amplification reaction occurs for one primer in the 5' Direction and for the other primer in the 3' Direction and wherein sufficiently complementary means said oligonucleotide sequence comprises at least seven consecutive nucleotides homologous to such a Direct Repeat sequence in particular the Sequence provided in Table II and/or exhibits at least 60% homology, preferably at least 80% homology. most preferably more than 90% homology with the corresponding part of the Direct Repeat sequence. In particular such a pair comprises one primer DRa capable of elongation in the 5' Direction and the other primer DRb capable of elongation in the 3' Direction with DRa being complementary to a sequence of the Direct Repeat located to the 5' side of the sequence of the Direct Repeat to which DRb is complementary, the Direct Repeat being present in the Direct Region of S. typhimurium.

Kits for carrying out a differentiation method according to any

of the described embodiments also fall within the scope of the invention. Such kits comprise a primer pair according to any of the described embodiments and optionally an oligonucleotide probe or a carrier, said carrier comprising at least 1 oligonucleotide probe specific for a spacer region of a bacterium to be determined said bacterium not belonging to ${\tt M}$ tuberculosis complex, preferably the oligonucleotide probe as defined, being an oligonucleocide probe of at least 10 nucleotides, preferably more than 12 nucleotides, in particular comprising between 12 to 40 nucleotides, said probe being sufficiently homologous to any of the spacer sequences or to fragments or derivatives of such spacer sequences to hybridise to such a spacer sequence, said oligonucleotide probe comprising at least 10 consecutive nucleotides homologous to such a spacer sequence and/or exhibiting at least 60% homology, preferably exhibiting at least 80% homology, most preferably exhibiting more than 90% homology with the corresponding part of the spacer sequence. Suitably a kit according to the invention comprises a data carrier with required reference patterns of the bacterial strain to be determined.

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DESCRIPTION OF THE FIGURES

Figure 1 depicts the structure of the DR region of M. bovis BCG as determined previously by Hermans et al. and Groenen et al. (12, 15). For the sake of convenience we will designate a DR plus its 3'adjacent spacer sequence as a "Direct Variant Repeat" (DVR). Thus, the DR region is composed of a discrete number of DVR's, each consisting of a constant part (DR) and a variable part (the spacer).

Figure 2 depicts multiple, short synthetic oligomeric DNA sequences based on the sequences of the unique spacer DNA's within the DR region.

Figure 3 shows the genetic organisation of the structures of the DR-like loci in various bacterial species.

15 be depicts the transcription direction of open reading frame (ORF)

For M. tuberculosis: MTCY 16B7.26, 27 and 30C are unknown genes/proteins.

For E. coli: iap gene function is alkaline phosphatase isozyme conversion. ORF f94, f305, YGCE and f223 are unknown genes/proteins.

For S. pyogenes: ORF1 and 2 are unknown genes/proteins.

For T. thermophilus: ORFC and D are unknown genes/proteins and ORF 1A and 1B are possibly transposases of IS elements 1000 and 1000A.

For Anabaena: No ORFs were annotated in the flanking sequences. The 130 bp insert is of unknown origin.

For Haloferax mediterranei: ORF21 is an unknown gene/protein. Probably another repeat cluster is also present on the megaplasmid pHM500.

For Methanococcus jannaschii: Comprises about 10 repeat clusters, the largest one of which comprises 25 repeats. All repeat clusters are coupled to a Long Repeat (LR) segment of 425bp. There are 18 LR's, some of which contain only one repeat. Smaller LR segments are also present, Δ LR. In one case, a cluster contains 5 repeats without LR (see ref. 18)

For M. thermoautrophicum: Two repeat clusters SRI and SRII flanked by

LRI, LRII. LRI and LRII are almost identical and are homologues of the LR segment of M. jannaschii. SRI and SRII are separated by 500 kb in the genome.

- For Thermatoga maritima: CelA gene encodes cellulase: endo-1.4-beta-glucanase (EC 3.2.1.4) and CelB is also a cellulase exhibiting 58% identity with celA.
- For Archaeoglobus fulgidus: The SRIA and SRIB repeat clusters have the same Repeat Sequence and the SRII Repeat Sequence is also clearly homologous. The SR clusters are separated by about 400bp. SRIB and SRII are located near tRNA genes. SRIA lies adjacent to an unknown ORF3.

Figure 4

- Hybridization Patterns of 17 E. coli isolates. Thirty four different spacer oligonucleotides were covalently linked to a membrane and PCR amplified DNA of E. coli was hybridized as described (Kamerbeek et al. 1997), except that the primers used to amplify the DR locus were specific for the DR sequence from E. coli. Note the polymorphism observed in E. coli due to the strain-dependent presence or absence of spacer DNA.
 - Figure 5

Hybridization Patterns of 4 Salmonella typhimurium isolates. Six different spacer oligonucleotides were covalently linked to a membrane and PCR amplified Salmonella DNA was hybridized as described (Kamerbeek et al 1997), except that the priners used to amplify the DR locus were specific for the DR locus of E. coli. Note the polymorphism observed in Salmonella due to the strain-dependent presence or absence of spacer DNA.

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M.bovis BCG

GICGICAGACCCCAAAACCCCGAGAGGGGGACGGAAAC Consensus DR sequence:

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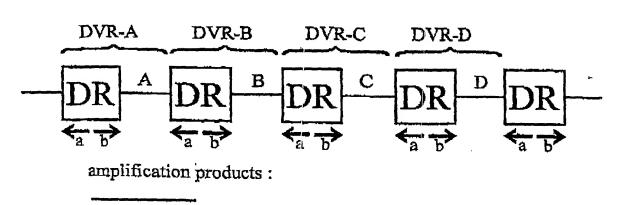
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Fig 2

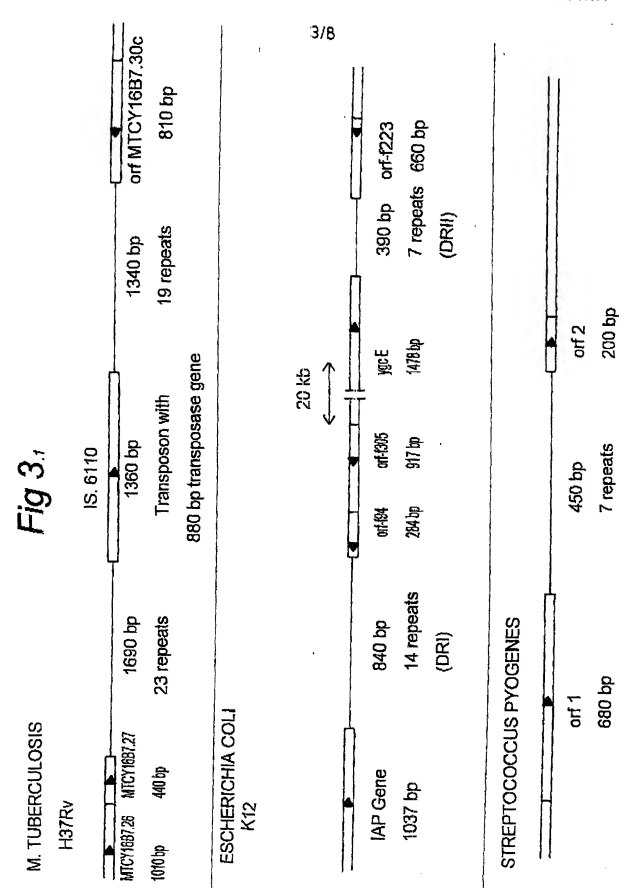


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4/8 180 bp 3 repeats orf 122.1 (380 pb) ORF 1b (960 bp) 8 repeats (540 bb) IS 1000 4 repeats 250 bp 21 repeats (1500 bp) Fig 3.2 130 bp orf 1A (960 bp) IS 1000A 9 repeats THERMUS AQUATICUS THERMOPHILUS 600 bp 4 repeats 250 bp HALOFERAX MEDITERRANE! ANABAENA sp. PCC 7120 720 bp 970 bp HB8 ATCC 27634 orf D orf C

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Na atherra duants bus 115

450 bp

3071 bp

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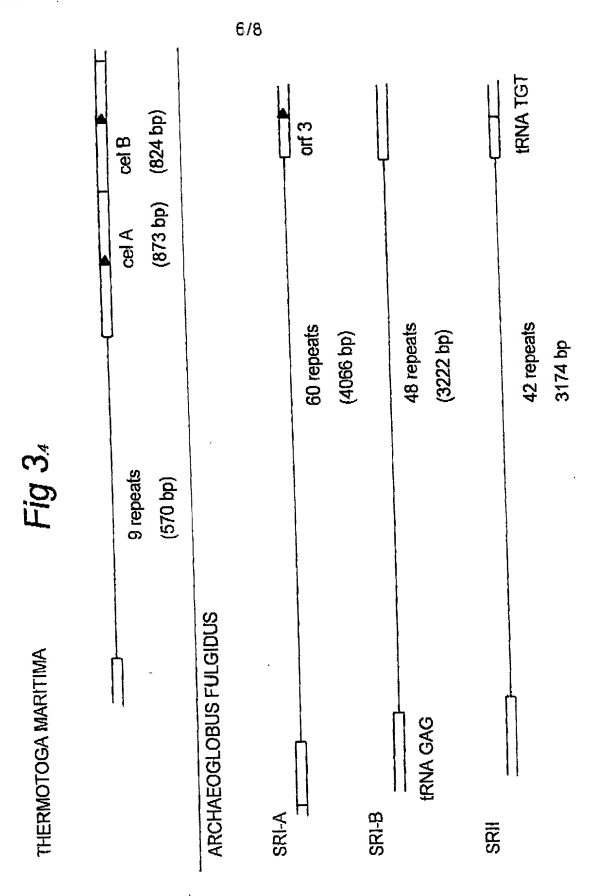
5/8 distance **√**500 kb 384 bp 2 orf M J 0001 (1127 bp) **MTH 1605** 5 repeats (310 bp) 47 repeats 25 repeats (1690 bp) Fig 3.3 125 repeats (8211 bp) METHANOBACTERIUM THERMOAUTOTROPHICUM 2 repeats METHANOCOCCUS JANNASCHII (30pb), MTH1075 (dq 099) LR-segment

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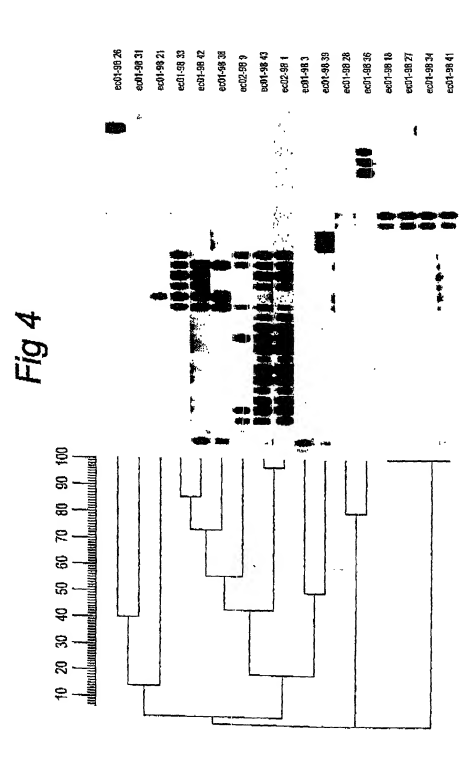


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Fig. 6

SEQ ID No. 1: GTCGTCAGACCCAAAACCCCGAGAGGGGACGGAAAC SEQ ID No. 2: CGGTTTATCCCCGCTGGCGCGGGAACTC SEQ ID No. 3: CGGTTTATCCCCGCTGGCGCGGGGAACTC SEQ ID No. 4: CGGTTTATCCCCGCTGGCGCGGGGAACTC SEQ ID No. 5: CGGTTTATCCCCGCTGGCGCGGGGAACTC SEQ ID No. 6: CGGTTTATCCCCGCTGGCGCGGGGAACTC SEQ ID No. 7: CGGTTTATCCCCGCTGGCGCGGGGAACTC SEQ ID No. 8: CGGTTTATCCCCGCTGGCGCGGATACAC SEQ ID No. 9: GTTTTAGAGCTATGCTGTTTTGAATGGTCCCAAAACT SEQ ID No. 10: AATCCCCTTACGGGGCTCAATCCCTTGCAA SEQ ID No. 11: GTTTCAATACTTCCTTAGAGGTATGGAAAC SEQ ID No. 12: GTTTTAACTAACAAAAATCCCTATCAGGGATTGAAAC SEQ ID No. 13: GTTTAAACTTTATAAAATCCCTTTTAGGGATTGAAAC SEQ ID No. 14: GTTACAGACGAACCCTAGTTGGGTTGAAGC SEQ ID No. 15: AATTAAAATCAGACCGTTTCGGAATGGAAA SEQ ID No. 16: ATTTCAATCCCATTTTGGTCTGATTTTAAC SEQ ID No. 17: GTTAAAATCAGACCAAAATGGGATTGAAAT

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SEQ ID No. 18: CTTTCAATCCCATTTTGGTCTGATTTCAAC

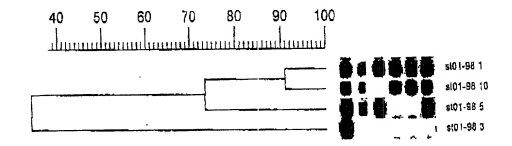
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Fig 5



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Fig. 6

SEQ ID No. 1: GTCGTCAGACCCAAAACCCCGAGAGGGGACGGAAAC

SEQ ID No. 2: CGGTTTATCCCCGCTGGCGCGCGCAACTC

SEQ ID No. 3: CGGTTTATCCCCGCTGGCGCGGGGAACTC

SEQ ID No. 4: CGGTTTATCCCCGCTGGCGCGGGGAACTC

SEQ ID No. 5: CGGTTTATCCCCGCTGGCGCGGGGAACTC

SEQ ID No. 6: CGGTTTATCCCCGCTGGCGGGGGAACTC

SEQ ID No. 7: CGGTTTATCCCCGCTGGCGCGGGGAACTC

SEQ ID No. 8: CGGTTTATCCCCGCTGGCGCGGATACAC

SEQ ID No. 9: GTTTTAGAGCTATGCTGTTTTGAATGGTCCCAAAACT

SEQ ID No. 10: AATCCCCTTACGGGGCTCAATCCCTTGCAA

SEQ ID No. 11: GTTTCAATACTTCCTTAGAGGTATGGAAAC

SEQ ID No. 12: GTTTTAACTAACAAAAATCCCTATCAGGGATTGAAAC

SEQ ID No. 13: GTTTAAACTTTATAAAATCCCTTTTAGGGATTGAAAC

SEQ ID No. 14: GTTACAGACGAACCCTAGTTGGGTTGAAGC

SEQ ID No. 15: AATTAAAATCAGACCGTTTCGGAATGGAAA

SEQ ID No. 16: ATTTCAATCCCATTTTGGTCTGATTTTAAC

SEQ 12 17: GITAAAATCAGACCAAAATGGGATTGAAAT-

SEQ ID No. 78: CTTTCAATCCCATTTTGGTCTGATTTCAAC

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(ORIGINAL DESIGN, NATIONAL STAGE OF PCT OR CIP APPLICATION)

As a below named inventor, I hereby declare that

My residence, post office address and citizenship are as stated below next to my name, I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled:

A method of interstrain differentiation of bacteria

the specification of which: (complete (a), (b) or (c) for type of application)

REGULAR OR DESIGN APPLICATION

as Application
and was amended on

PCT FILED APPLICATION ENTERING NATIONAL STAGE

c. [X] was described and claimed in International application No. PCT/NL98/00186 filed on 3 April 1998 and as amended on (if any)

ACKNOWLEDGEMENT OF REVIEW OF PAPERS AND DUTY OF CANDOR

I hereby state that I have reviewed and understand the contents of the above identified specification, including the claims, as amended by any amendment referred to above.

I acknowledge the duty to disclose information which is material to patentability as defined in Title 37, Code of Federal Regulations, paragraph 1.56(a).

In compliance with this duty there is attached an information disclosure statement 37 CFR 1.97

PRIORITY CLAIM

I hereby claim foreign priority benefits under Title 35. United States Code paragraph 119 of any foreign application (s) for patent of inventor's certificate listed below and have also identified below any foreign application for patent of inventor's certificate having a filing date before that of the application on which priority is claimed.

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(complete (d) or (e))

d. [] no such applications have been filed such applications have been filed as follows e.[]

EARLIEST FOREIGN APPLICATION(S), IF ANY FILED WITHIN 12 MONTHS (6 MONTHS FOR DESIGN) PRIOR TO SAID APPLICATION

Country	Application Number	Date of filing (day, month, year)	Date of Issue (day, month, year)	Priority claimed

ALL FOREIGN APPLICATION(S), IF ANY FILED MORE THAN 12 MONTHS	S
(6 MONTHS FOR DESIGN) PRIOR TO SAID APPLICATION	

CONTINUATION-IN-PART

(Complete this part only if this is a continuation-in-part application)

I hereby declare claim the benefit under Title 35, United States code, paragraph 120 of any United States application(s) listed below and, insofar as the subject matter of each of the claim of this application is not disclosed in the prior United States application in the manner provided by the first paragraph of Title 35, United States Code, paragraph 112, I acknowledge the duty to disclose material information as defined in Title 37, Code of Federal Regulations, paragraph 1.56(a) which occurred between the filing date of the prior application and the national or PCT international filing date of this application;

٠.	(Application Serial No.)	(Filing date)	(Status)	(patented, pending, abandoned)
	(Application Serial No.)	(Filing date)	(Status)	(patented, pending, abandoned)

POWER OF ATTORNEY

As a named inventor, I hereby appoint the following attorney(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith: Robert J. PATCH, Reg. No. <u>17,355.</u> Andrew J. PATCH, Reg. No. 32,925, Robert F. HARGEST, Reg. No. 25,590, Benoît CASTEL, Reg. No. 35,041, Eric Jensen, Reg. No. 37,855, and Thomas W. PERKINS, Reg. No. 33,027 c/o YOUNG & THOMPSON, Second Floor, 745 South 23rd Street, Arlington, Virginia 22202.

Address all telephone calls to Young & Thompson at 703/521-2297.

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